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File No: 135555-0262

Box PATENT APPLICATION

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S I R :

Enclosed please find a continuation application for United States patent as identified below:

Inventor/s (name ALL inventors): Douglas W. Walker, Frank M. Ordaz,  
and M. Terry Olson.

Title: CORDLESS SURGICAL HANDPIECE WITH DISPOSABLE BATTERY; AND  
METHOD

including the items indicated:

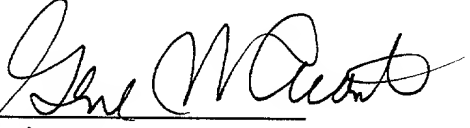
1. Specification and 28 claims: 10 indep.; 18 dep.
2. [X ] Declaration and power of attorney
3. [X] Formal drawings, 7 sheets (Figs. 1-13)
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5. This application claims the benefit of:  
Application SN 09/349,643 filed 7/8/99 and  
Provisional Application 60/112,678 filed 12/16/98

Respectfully submitted,

Dated: September 22, 2000

  
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**File 153635-0262**

United States Patent Application

for

CORDLESS SURGICAL HANDPIECE WITH DISPOSABLE BATTERY; AND METHOD

Inventors: Douglas W. Walker,  
Frank Ordaz, and M. Terry Olson

Claiming the Benefit of  
Application S. N. 09/349,643 filed 07/08/99  
and  
Prov. Appln. 60/112,678 filed Dec. 16, 1998

Assigned to Medtronic, Inc.

1 CORDLESS SURGICAL HANDPIECE WITH DISPOSABLE BATTERY; AND METHOD

2  
3  
4 PRIORITY CLAIM  
5  
6

7 This application claims the benefit of our prior copending  
8 application S. N. 09/349,643 filed 07/08/99 and of our  
9 Provisional Application 60/112,678 filed Dec. 16, 1998  
10

11 FIELD OF THE INVENTION

12 The present invention relates to electrically operated  
13 surgical tools and methods of their use.  
14

15 BACKGROUND OF THE INVENTION

16 An important economic factor for any surgical tool or  
17 machine is the amount of unproductive time spent in its  
18 operation. This comes in the form of training and initial set-  
19 up. Before a surgical procedure begins, the users need to know  
20 how to operate the instrument and be familiar with all its  
21 controls. Hospital personnel must set up the instrument system  
22 before it can be used. This involves connecting all the power  
23 lines, calibration, and verifying function. Both user's training  
24 and instrument system set-up entail time and cost by the medical  
25 facility.

26 The most unique aspect of surgical equipment is the need for  
27 sterility. To prevent infections and to aid healing, surgical  
28 equipment is sterilized. The effort this takes and the success

1 with which it occurs varies depending on the equipment. The  
2 maintenance of sterility in the surgical site is a major factor  
3 in health care. Non-sterile or partially sterile instruments may  
4 be considered to harbor bacteria or toxic debris. Each of these  
5 will lead to an inflammatory response from the body and  
6 associated infection, carcinogenesis and cell necrosis.  
7 Contamination will lead to grave systemic effects on the patients  
8 of orthopedic surgery, which by its nature is highly invasive.  
9 The maintenance of a sterile operating field is of prime  
10 importance in surgical handpieces.

#### 11 CORDED INSTRUMENT SYSTEMS

12 The use of a corded instrument system dictates that only one  
13 handpiece may be used at any one time per console. When another  
14 instrument is needed the console cord must be switched to another  
15 instrument. Procedures where it is imperative that two  
16 instruments be used simultaneously necessitate two consoles.  
17 Handpiece instrument consoles are expensive and the need for  
18 duplication is a distinct disadvantage. Disconnecting and  
19 reconnecting instruments to the console cord takes a certain  
20 amount of work and time. Using cordless instruments reduces  
21 instrument transfer time.

22 A reusable cord must be properly sterilized prior to  
23 surgery in an expensive and complex in-house sterilizer. Once  
24 sterile, the cord must be carefully transported to the operating  
25 room. There the cord becomes partially non-sterile due to its  
26 console connection.

1       When a corded handpiece is transferred and exchanged to  
2       other medical personnel, the cord must be handled. However,  
3       there is no way to tell which part of cord has fallen out of the  
4       sterile field. This poses an increased risk of contamination to  
5       the user and the surgical site.

6       When multiple handpieces are exchanged they are extensively  
7       handled. This is because the cord must be disconnected and  
8       reconnected to each handpiece. Handling increases the risk of  
9       contamination. This risk is greatly minimized with cordless  
10      instruments.

11      Following a surgical procedure, the instrument cord needs to  
12      be thoroughly cleaned prior to its next use. This involves  
13      cleaning blood and tissue off the cord with powerful solvents and  
14      cleaners. These agents attack the cord and limit its useful  
15      life. The cord must be handled by trained medical personnel.  
16      A disposable power pack would eliminate these extra tasks and  
17      their associated costs.

#### 19                                   CORDLESS INSTRUMENTS

20      During a surgical procedure it is often necessary to use  
21      more than one handpiece. Saws and high speed handpieces are  
22      used for cutting, shaping and in general removing portions of  
23      bone. Drills are used primarily for making holes, which are  
24      then used for inserting wires, pins or screws. These two  
25      operations are often used in conjunction with one another during  
26      bone and tissue repair procedures. Current pencil grip surgical

1 instruments make it difficult to use more than one instrument at  
2 a time during a surgical procedure.

3 A cordless handpiece is easier for the physician to operate  
4 than similar corded handpieces. The controls are basic and are  
5 controlled with one hand. Equivalent corded handpieces require  
6 complex console instructions and commands.

7 A further advantage of cordless handpieces is the greater  
8 ease of set-up. Equipment set-up is a significant time issue for  
9 hospitals. Corded systems have complicated and time-consuming  
10 assembly procedures. Multiple connections are involved. A user  
11 must be trained at setting up and operating the console system.

12 In instances where a powered surgical instrument is needed  
13 immediately, a cordless instrument can be immediately transported  
14 to that area. This is true regardless of the surrounding  
15 environment. That is not true of a corded unit.

16 A surgical instrument cord, connected to a non-sterile  
17 console, may be considered only partially sterile. This by  
18 itself compromises the integrity of a surgical site. A cordless  
19 handpiece maintains sterility throughout the entire surgical  
20 operation.

21 Surgical procedures often involve cuts at more than one  
22 position at a surgical site. A convenient way to accomplish this  
23 is to pass the surgical instrument over the surgical site. With  
24 a corded instrument, this would unfortunately result in the cord  
25 being passed over, and perhaps falling into, the surgical site.  
26 The resulting tissue damage and contamination can have grave

1 consequences. A cordless instrument can easily be passed over  
2 the surgical site without contamination risks. Likewise, a  
3 physician who needs to make multiple cuts or holes at various  
4 surgical locations needs to be in the position that best suited  
5 him or her without trailing a bulky, cumbersome partially sterile  
6 cord.

7 Cordless instruments have heretofore utilized batteries of  
8 the rechargeable type. This adds special problems, because the  
9 battery must be sterilized before use; and after each use it must  
10 be recharged and then sterilized in preparation for the next use.  
11 Presently available batteries do not lend themselves well to this  
12 process.

#### 13 SUMMARY OF THE INVENTION

14 According to the present invention a method of performing  
15 surgery is provided, in which bone or hard tissue may be cut,  
16 shaped, or drilled by means of a cordless powered surgical  
17 instrument, but without the necessity of subsequently recharging  
18 a battery or re-establishing its sterile condition.

19 According to the invention a disposable battery pack can be  
20 easily connected to a surgical instrument, both electrically and  
21 mechanically, and after a single use may be detached and safely  
22 disposed of as non-hazardous waste, into the waste system.

23 Further according to the invention a method of performing a  
24 surgical procedure is disclosed, utilizing a cordless surgical  
25 handpiece powered from a sterile battery pack in which the  
26



1 battery chemistry is based upon lithium/manganese dioxide, the  
2 battery being in condition for immediate use without further  
3 charging or sterilization, and being adapted after a single use  
4 to be disposed of into non-hazardous waste.

5 Still further according to the present invention a surgical  
6 handpiece and disposable battery are provided with mating sets of  
7 electrical contacts which can be mechanically and conductively  
8 locked together to ensure correct alignment of the parts, as well  
9 as stable mechanical support and reliable electrical operation of  
10 the handpiece during the surgical procedure.

11 According to yet another feature of the presently preferred  
12 form of the invention, a surgical handpiece and a disposable  
13 battery each has a defined longitudinal axis with a set of  
14 electrical contact elements arranged generally concentric to that  
15 axis, and when those parts are aligned on a mutual longitudinal  
16 axis the sets of contacts are adapted to become lockingly and  
17 conductively interengaged in response to rotation of the battery  
18 pack relative to the handpiece.

19 Still another feature of the invention is that when using a  
20 compact surgical handpiece with a brushless DC motor and a  
21 manually operated external trigger for activating and controlling  
22 the motor operations, a compatible disposable battery may also be  
23 used, with interengaging sets of contacts on the handpiece and  
24 battery adapted to become lockingly and conductively interengaged  
25 upon rotation of the battery pack relative to the handpiece in a  
26 manner that rapidly achieves correct alignment of the parts and

1 also ensures stable mechanical attachment and support during the  
2 surgical procedure.

### 3 DRAWING SUMMARY

4 Fig. 1 (a) is a schematic view of a surgical handpiece in  
5 accordance with the invention, showing disposal of an associated  
6 battery pack into non-hazardous waste after it has been used;

7 Fig. 1 (b) is a cross-section view taken on the Line 1(b) --  
8 (1 (b) of Fig. 1 (a), showing that the battery pack when being  
9 disposed of still contains batteries;

10 Fig. 2 is a perspective view of a sterile battery pack in  
11 its plastic container, in accordance with the invention;

12 Fig. 3 is an exploded perspective view of the handpiece and  
13 battery pack before they are assembled together;

14 Fig. 4 is a perspective view of the instrument assembly  
15 during attachment of the battery pack, with arrows indicating the  
16 direction of movement of the battery pack;

17 Fig. 5 is a perspective view of the instrument assembly  
18 after attachment of the battery pack, with arrows showing that  
19 the battery pack has been rotated by ninety degrees;

20 Fig. 6 is a rear end view of a surgical drill or handpiece  
21 in accordance with the invention taken along line 6-6 of Fig. 3;

22 Fig. 7 is a front end view taken along line 7-7 of Fig. 3 of  
23 the disposable battery pack;

24 Fig. 8 is a longitudinal cross-sectional view showing  
25 initial alignment of the battery pack to the surgical drill;

26 Fig. 9 is a cross-sectional view like Fig. 8, but showing

1 the battery pack after its full insertion and its rotation into a  
2 locking attachment;

3 Fig. 10 is a perspective view of the battery pack showing in  
4 detail its forward end;

5 Fig. 11 is a perspective view of the batteries contained  
6 within the battery pack;

7 Fig. 12 is a side elevation view of the battery pack; and

8 Fig. 13 is a perspective view of a complete instrument  
9 showing an alternate form of trigger for handpiece control.

#### 11 DETAILED DESCRIPTION

12 In Fig. 1(a) and 1 (b) a surgical handpiece 10 with housing  
13 12 has a rearward or base end 14, an externally mounted trigger  
14 16, and a tool 18 on its forward end. A disposable battery pack  
15 20 has a housing 22 and contains internal batteries 24. Arrow 30  
16 indicates the disposal of battery pack into a waste basket 32  
17 after the surgical handpiece has been used in a surgical  
18 procedure.

19 As is conventional, the surgical handpiece is a compact  
20 device containing a brushless DC motor for moving the tool member  
21 18, a manually operated trigger 16 used for activating motor  
22 control operations, and an adjustable tool support mechanism for  
23 orienting the tool 18 and securing it in place. An electronic  
24 circuit, not specifically shown, controls and regulates the  
25 energy supply to the motor, and is operated externally by the  
26 trigger 16. Trigger 16 may also control the range of speed, the

1 direction of the cutting tool, and cutting tool braking.

2 According to the present invention the energy supply for the  
3 handpiece is provided by the battery pack 20 which is secured  
4 onto the handpiece by means of a rotating movement. This  
5 detachable DC electric energy supply ensures correct orientation  
6 of the electrical contacts in the battery to those for the  
7 handpiece, and also signals to the operator by sound, sight, and  
8 touch that the battery is correctly secured in place.

9 The battery pack 20 is pre-sterilized and packaged for  
10 immediate use in a surgical environment. Fig. 2 shows a plastic  
11 cover 26 which is preferably used to enclose the battery pack  
12 prior to its use. The battery pack 20 consists of primary  
13 batteries 24 which by definition do not require charging before  
14 use. These batteries retain their initial charge for long  
15 periods of time. The battery pack is discarded after use and may  
16 be considered a disposable component of the instrument system.

17 Further according to the invention the battery pack  
18 preferably contains primary batteries 24 whose chemistry is based  
19 upon lithium/manganese dioxide, such as the DL 2/3A manufactured  
20 by Duracell, Inc. of Bethel, Connecticut. These batteries  
21 possess a high energy density, have a high rate capability over a  
22 broad temperature range, and have excellent capacity retention.

23 As shown in Fig. 3 the handpiece 10 at its rearward or  
24 battery receiving end 14 has an alignment post 40 extending  
25 therefrom, which defines a longitudinal axis of the handpiece.  
26 It also has a set of electrical contact elements 42a and 42b

1 which are concentric to that axis. The sterile and disposable  
2 battery pack 20 has an attachment end 50 with a central opening  
3 52 therein, the opening 52 also defining a longitudinal axis of  
4 the battery pack. The battery pack 20 also has a set of  
5 electrical contact elements 54 which are concentric to its  
6 longitudinal axis. The central opening 52 in the disposable  
7 battery pack 20 is adapted to insertably receive the alignment  
8 post 40 so as to establish a mutual alignment axis of the  
9 handpiece 10 and battery pack 20.

10 The battery receiving end of the handpiece 10 also has  
11 flat end surface surfaces 44a and 44b which are adapted to be  
12 engaged by the battery pack. The forward or attachment end of  
13 the disposable battery pack has a flat end surface 56 adapted for  
14 abutting engagement with the end surfaces 44a and 44b while yet  
15 allowing relative rotation of the battery pack relative to the  
16 handpiece.

17 The set of contacts 42a and 42b on the battery receiving end  
18 of the handpiece and the set of contacts 54 on the attachment end  
19 of the disposable battery pack are mating sets of electrical  
20 contact elements, each set being arranged generally concentric to  
21 the mutual alignment axis. Upon the insertion of the alignment  
22 post 40 of the handpiece into the opening 52 of the battery pack,  
23 the sets of mating contacts are adapted to then become lockingly  
24 and conductively interengaged in response to rotation of the  
25 battery pack relative to the handpiece.

26 In operation, the post 40 is first partially inserted into

1 opening 52 to establish alignment of battery and handpiece. Then  
2 with further insertion, the two sets of contacts will assume a  
3 position in concentric relation to their common mutual axis of  
4 alignment.

#### 7 THE INTERLOCKING PARTS

8 Referring now to Figs. 3, 6, 8, 9, and 12, the detailed  
9 structure of the rearward end of handpiece 12 is shown. Rearward  
10 end 14 of the housing of handpiece 12 has an aluminum cover. An  
11 end plate 60 made of aluminum closes the rearward end of  
12 handpiece housing, as best seen in the cross-section views of  
13 Figs. 8 and 9. End plate 60 is recessed inwardly from the  
14 extreme rearward end of the housing.

15 There are a pair of electrical contacts 42a and 42b which  
16 protrude out from end plate 60. Those contacts do not support  
17 themselves, however; a plastic cylinder 62 is secured to end  
18 plate 60, and the contacts 42a 42b, are secured to the outer wall  
19 of plastic cylinder 62, about 180 degrees apart. One contact is  
20 of course positive, and the other negative.

21 The rearward end of the handpiece also has flanges that are  
22 part of and protrude outward from end plate 60 to control the  
23 insertion and locking of the battery pack, designated on Fig. 3  
24 by numerals 44a and 44b. On the left side as seen in Fig. 3,  
25 **there is a wide** gap between 44a and 44b. On the right side as  
26 seen in Fig. 3, there is a **narrow** gap. The flanges are

1 preferably formed as an integral part of the end plate 60, as  
2 shown in Fig. 8. A stop pin 65 seen in the upper right portion  
3 of Fig. 6 protrudes inwardly behind flange 44b and limits the  
4 rotation of the battery contacts relative to the handpiece.

5 The front end of battery pack 20 is shown in Figs. 3, 7, 8,  
6 9, and 10. It has a front end plate 70, formed of plastic  
7 material, such as ABS plastic; see Fig. 8. End plate 70 also has  
8 a projecting ring 72, with flanges 74, 76. As best seen in Fig.  
9 3, flange 74 on the near side of the battery pack will fit into  
10 the gap between lower flange 44a and upper flange 44b of the  
11 handpiece, which gap is also on the left as seen in Fig. 3.

12 The other flange 76 on far side of battery pack will fit  
13 between flanges 44a, 44b on the far or right side of the  
14 handpiece as seen in Fig. 3. However, flange 74 is **too wide** to  
15 enter the gap on the right side as seen in Fig. 3. Therefore,  
16 battery pack 20 must be engaged with the handpiece in a  
17 predetermined relative position.

18 Fig. 8 shows alignment of the two parts of apparatus on  
19 their mutual longitudinal axis as the post 40 makes its initial  
20 entry into the center hole 52 of the battery pack. Further  
21 insertion of the post ensures the coaxial alignment of the two  
22 parts.

#### 23 24 ROTATING THE BATTERY PACK TO LOCKED POSITION

25 Then battery pack 20 is rotated to the right, as indicated  
26 by arrows 80 in Figs. 3, 4, and 5. Fig. 5 shows the locked

1 position, also shown in more detail in Fig. 9. Further rotation  
2 of the battery pack relative to the handpiece is prevented by the  
3 stop pin 65. The contact elements 42a and 42b are made as spring  
4 members, so that when the contact elements 54 of the battery pack  
5 are seated, there is an audible noise to tell the operator that  
6 the properly aligned operating position has been reached. Thus  
7 the apparatus includes means providing a spring-supported snap  
8 action so that the sets of mating contacts become lockingly and  
9 conductively interengaged in response to rotation of the battery  
10 pack relative to the handpiece.

11 It will also be noted that the battery receiving end of the  
12 handpiece, and the attachment end of the battery pack, each has a  
13 non-circular external cross-sectional configuration. Thus in the  
14 presently preferred embodiment of the invention the two housings  
15 are essentially square with rounded corners. The two external  
16 configurations are closely similar in both size and shape, and  
17 the rotational position of the battery pack when the contacts are  
18 locked together is such that the handpiece and the battery pack  
19 then provide an essentially continuous external surface. This  
20 indicates to the hand of the operator that correct alignment of  
21 the contacts has been achieved.

#### 2 ADVANTAGES OF CORDLESS SURGICAL HANDPIECES

3 Orthopedic surgical instruments are widely used in many  
4 delicate bone working procedures. These include spinal  
5 surgeries, neurosurgeries and other bone sculpting operations.  
6 For these surgeries, a small, light weight, well balanced



1 instrument is desired. The handpiece needs to have high speed  
2 and power, and be sterilizable.

3 The ergonomics of a pencil grip cordless instrument is  
4 quantifiably better than corded or pistol grip handpieces.  
5 Pencil grip allows more, fine motor control and easier eye-hand  
6 coordination. The improved ergonomics of the pencil grip results  
7 in less fatigue by the user.

8 A cordless handpiece is more mobile than a corded unit.  
9 This makes it easier to pass it around the surgical site, from  
10 physician to attendant or assistant. A cordless system is also  
11 not weighed down by a cord. The back of corded or pistol grip  
12 handpieces can require tugging or lifting of the handpiece  
13 cutting tool, which makes cutting more difficult. The cord  
14 diameter is very thick for a high performance system, which again  
15 limits mobility.

16 A further disadvantage of a corded instrument is that it is  
17 tethered. The handpiece can only be operated at a given distance  
18 from the console, restricted by the length of the cord. The cord  
19 also has associated inductance and capacitance properties which  
20 can affect the handpiece performance or electrical complexity of  
21 the system.

22 For movement about the surgical site with a corded  
23 handpiece, extra cord is needed. The extra cord is either coiled  
24 by the physician, held by an attendant or attached to a surgical  
25 stand. This coiling necessitates extra labor by operating room  
26 (OR) personnel, takes up critical surgical site space and limits  
27 mobility of other OR personnel.

28 The mobility and transport of cordless, pencil grip  
29 instruments is of particular advantage when multiple instruments

1 are needed in a procedure. This is often the case where  
2 drilling, sawing and wire driving are all needed. These often  
3 come in rapid succession and in different sequences. Time is a  
4 critical factor for successful surgical outcomes.

#### 5 ELECTRICAL ADVANTAGES OF CORDLESS INSTRUMENTS

6 Instrument performance is of critical importance in surgical  
7 procedures. A key electrical factor with corded handpieces is  
8 the cord itself. The power cord has a given length associated  
9 with it and electrical properties of its own.

10 The internal impedance of the cord increases electrical  
11 losses and reduces power and performance. The power is lost as  
12 heat and many have other electro - magnetic interference (EMI)  
13 problems associated with it. The length of cord and flexibility  
14 needed usually necessitate the use of three-phase current.

15 The cord also has associated inductance and capacitance  
16 properties which reduces performance and necessitates more  
17 complex electronics in the console. There are also associated  
18 EMI problems with this inductance and impedance. These factors  
19 exist with both DC and, especially, AC signals.

20 The electrical properties of the cord are directly  
21 proportional to its length and can only be minimized by  
22 shortening the cord or making it thicker. A shorter cord reduces  
23 mobility. A thicker cord reduces flexibility and increases  
24 costs.

25 The EMI problems associated with a cord occur along the  
26 entire cord length. The two connections are especially prone to  
27 emissions. Cordless instruments, on the other hand, run on  
28 direct current (DC), have only one connection point and minimized  
29 electronics. The basic level of emitted EMI from the battery to

1 handpiece connection, in a cordless handpiece, is zero.

2 A corded surgical instrument has a direct connection between  
3 the patient and wall or line voltage. Power surges from the  
4 outlet must be properly controlled before they reach the patient.  
5 A cordless instrument which operates with lower voltage, less  
6 energy, and a limited capacity power supply, poses a much lower  
7 electrical threat to the patient.

8 The calibration and preventive maintenance costs associated  
9 with a corded handpiece console are not a factor with a cordless  
10 system. The elimination of the console results in more operating  
11 room table space. This valuable space can then be used for  
12 important instruments and equipment which need to be close at  
13 hand.

#### 14 15 COST AND RELIABILITY OF CORDLESS SYSTEMS

16 The cost and reliability of any surgical instrument is of  
17 critical importance. OR expenses are high and are based on time  
18 usage. All general surgeries have a time factor based on how  
19 long the room is occupied and the patient is under anesthesia.  
20 Instrument systems must properly work or the operation's success,  
21 and thus the patient's health, will be affected.

22 Cord breakage or damage is a prevalent problem and  
23 necessitates cord replacement or repair. Cord failure may also  
24 damage the handpiece and/or console through electrical shorts or  
25 otherwise. Even partial failure will directly decrease the  
26 performance of the surgical system. Cord repair and replacement  
27 is expensive, as is system repair and replacement.

28 Due to the high failure rate of cords, as well as their  
29 importance to the handpiece system, they must be tested often.

1 This results in time and expense to the medical facility, even  
2 when they are functional.

3 Cord damage may result in heat build-up or exposed  
4 electrical leads. These pose serious risks to the operators as  
5 well as the patient.

6 Sterilization and cleaning using hospital grade detergents  
7 have serious negative effects on the cord. These thermal and  
8 chemical agents directly reduce the life of the cord, as well as  
9 increase the likelihood of damage.

10 Cord damage may also result in improper system  
11 communications and linkage. This may result in increased cord  
12 impedance and consequent loss of handpiece performance. Improper  
13 signaling may also lead to improper motor operations and result  
14 in handpiece overheating. This may be a hazard to either the  
15 user, patient or attending personnel.

#### 16 17 ADVANTAGES OF DISPOSABLE BATTERY PACKS

18 The high operating costs of surgical arenas and medical  
19 personnel dictate the use of low maintenance, easy to use  
20 equipment. Many times this means single use or disposable  
21 components. Also supporting disposable product use is the issue  
22 of sterility and contaminants from the surgical site. Surgical  
23 equipment is often made disposable whenever possible. Packaged  
24 pre-sterile, disposable medical equipment, by its nature, is easy  
25 to use and maintain and often very cost-effective.

26 Disposable, primary battery packs offer a higher energy  
27 capacity per volume than equivalent rechargeable batteries. The  
28 batteries are smaller, which is a distinct advantage in cordless  
29 surgical handpieces.

1 Disposable, primary battery packs offer a higher energy  
2 capacity per weight than equivalent rechargeable batteries. The  
3 batteries are lighter, which is a distinct advantage in cordless  
4 surgical handpieces.

5 Disposable batteries are inspected at the manufacturer by  
6 trained personnel prior to each use. This gives the products a  
7 higher reliability per use than equivalent reusable batteries.  
8 This is an advantage during critical use situations that often  
9 occur in the operating room.

10 Every battery pack has full running power immediately upon  
11 attachment. Rechargeable batteries, on the other hand, must be  
12 tested to verify their immediate charge capacity.

13 Due to the nature of disposable batteries there is no pre-  
14 or post operation clean-up involved. This reduces the amount of  
15 time hospital personnel need to service the equipment.

16 Reusable batteries will exhibit wear after repeated use.  
17 This results in debris build-up on components, particularly  
18 oxidation on electrical contacts, weakening of the housing and  
19 general degradation. This may result in poor or unacceptable  
20 product performance. Disposable products do not have these  
21 associated problems.

22 Disposable, primary batteries may be entered into the normal  
23 waste stream. Standard rechargeable batteries are considered  
24 toxic and must be properly disposed of outside the normal waste  
25 stream.

26 Most disposable primary batteries may be transported by air  
27 or other standard methods. Some types of rechargeable batteries  
28 need special handling conditions and permits to be safely  
29 transported.

1 By their very nature disposable primary battery cells have  
2 an immediate charge capacity, unlike rechargeable battery cells.  
3 Rechargeable batteries need to be charged on specialized and  
4 dedicated chargers. This is an expensive and bulky piece of  
5 equipment for the medical facility.

6 Rechargeable, secondary batteries need to be recharged  
7 immediately prior to each use because they lose their electrical  
8 charge very quickly. Primary batteries retain their charge for  
9 very long periods of time, over ten years. This results in lower  
10 service and maintenance costs to hospital for disposable  
11 batteries.

12 Primary batteries have a much lower cost base than  
13 equivalent rechargeable batteries. This usually has the  
14 advantage of saving money for medical facilities.

15 Disposable primary cells generally have a higher voltage,  
16 and thus a higher energy potential than rechargeable cells.  
17 Furthermore, a higher voltage allows most standard electrical and  
18 electromechanical components and motors to operate more  
19 efficiently, with fewer components, at a lower cost.

20 Disposable, primary batteries retain their charge capacity  
21 under more varied conditions, including thermal, humidity and  
22 mechanical (e.g. vibration or shock) than equivalent rechargeable  
23 battery cells. This increased robustness is an advantage in  
24 surgical operations.

25 In improper use conditions, such as a short circuit, many  
26 types of batteries will vent their internal electrolyte in the  
27 form of a gas. If, due to improper conditions, primary batteries  
28 vent internal gas, these gases are non-toxic. This is unlike  
29 some types of rechargeable batteries. That can be a critical

1 issue in an operating room where it would be very difficult to  
2 evacuate the area if this condition arose.

3  
4 ALTERNATE EMBODIMENT

5 In the alternate embodiment of the present invention as  
6 shown in Fig. 13, the handpiece 10' has a rather long-handled  
7 trigger 16'.

8 While the presently preferred embodiment of the invention  
9 has been disclosed in detail in order to comply with requirements  
10 of the patent laws, it will be understood by those skilled in the  
11 art that some variations may be possible within the concept of  
12 the invention. It will therefore be understood that the scope of  
13 the invention is to be determined only in accordance with the  
14 appended claims.

15 WHAT WE CLAIM IS:  
16

1  
1 1. A surgical instrument comprising:  
2 (a) a handpiece having a tool supporting end, and a battery  
3 receiving end;  
4 (b) a battery pack having an attachment end;  
5 (c) one of the battery receiving end and attachment end  
6 having an alignment post with a plurality of electrical contacts  
7 arranged concentric thereto, and the other thereof having a  
8 central opening defining a longitudinal axis with a set of  
9 electrical contact elements concentric to that axis; and  
10 (d) wherein the two sets of contacts are adapted to become  
11 lockingly and conductively interengaged upon engagement of the  
12 alignment post with the central opening and in response to  
13 rotation of the battery pack relative to the handpiece.

2. A surgical instrument as in Claim 1 wherein the battery  
pack has chemistry based upon lithium/manganese dioxide, the  
battery pack after use being disposable into non-hazardous waste.



1           3. A surgical instrument for performing a cutting, shaping,  
2 or drilling operation on bone or hard tissue, comprising:

3           (a) a handpiece having a tool supporting end, and a battery  
4 receiving end with a set of electrical contact elements thereon;

5           (b) a sterile package containing a disposable battery, the  
6 battery chemistry being based upon lithium/manganese dioxide;

7           (c) the disposable battery having an attachment end with a  
8 set of electrical contact elements on its attachment end;

9           (d) the handpiece and the battery each having a defined  
10 longitudinal axis, each set of electrical contact elements being  
11 arranged generally concentric to that axis, and wherein in  
12 response to rotation of the battery pack relative to the  
13 handpiece the sets of contacts are adapted to become lockingly  
14 and conductively interengaged prior to the surgical procedure.

1           4. A surgical instrument comprising:

2           (a) a handpiece having a tool supporting end, and a battery  
3 receiving end with an alignment post extending therefrom, the  
4 battery receiving end of the handpiece also having a set of  
5 electrical contact elements arranged in generally concentric  
6 relation to the alignment post;

7           (b) a battery pack having an attachment end with a central  
8 opening therein, and a set of mating electrical contact elements  
9 arranged in a generally circular configuration concentric to the  
10 central opening;

11           (c) the sets of mating contacts being adapted to come into a  
12 mutually concentric relation in response to insertion of the  
13 alignment post into the central opening;

14           (d) the sets of contacts upon rotation of the battery pack  
15 relative to the handpiece being adapted to then become lockingly  
16 and conductively interengaged in a predetermined relative  
17 position; and

18           (e) means indicating by at least one of sight, sound, and  
19 touch that the predetermined relative position has been achieved.

1           5. A surgical instrument for performing a cutting, shaping,  
2 or drilling operation on bone or hard tissue, comprising:

3           (a) a handpiece having a battery receiving end with an  
4 alignment post extending therefrom;

5           (b) a sterile package containing a disposable battery pack  
6 which has an attachment end with a central opening therein;

7           (c) the central opening in the disposable battery pack being  
8 adapted to insertably receive the alignment post so as to  
9 establish a mutual alignment axis of handpiece and battery pack;

10           (d) the battery receiving end of the handpiece and the  
11 attachment end of the disposable battery pack having flat end  
12 surfaces adapted for abutting engagement while yet allowing  
13 relative rotation of the battery pack relative to the handpiece;

14           (e) the battery receiving end of the handpiece and the  
15 attachment end of the disposable battery pack having mating sets  
16 of electrical contact elements, each set being arranged generally  
17 concentric to the mutual alignment axis; and

18           (f) wherein upon the insertion of the alignment post of the  
19 handpiece into the opening of the battery pack, the sets of  
20 mating contacts are adapted to then become lockingly and  
21 conductively interengaged in response to rotation of the battery  
22 pack relative to the handpiece.

6. The apparatus of Claim 5 wherein the chemistry of the  
disposable battery pack is based upon lithium/manganese dioxide.

7. The apparatus of Claim 5 including means providing a spring-supported snap action whereby the sets of mating contacts become lockingly and conductively interengaged in response to rotation of the battery pack relative to the handpiece.

8. The apparatus of Claim 7 wherein the spring-supported snap action means provides an audible sound indicating that the mating contacts and the battery have been correctly and securely locked in position.

9. The apparatus of Claim 5 wherein the battery receiving end of the handpiece, and the attachment end of the battery pack, each has a non-circular external cross-sectional configuration, the two external configurations being closely similar in both size and shape, and the rotational position of the battery pack for locking the contacts being such that the handpiece and the battery pack then provide an essentially continuous external surface which indicates to the hand of the operator that correct alignment of the contacts has been achieved.

10. The apparatus of Claim 9 including means providing a spring-supported snap action whereby the sets of mating contacts become lockingly and conductively interengaged in response to rotation of the battery pack relative to the handpiece.

1           11. A surgical instrument for performing a surgical  
2 procedure on bone or hard tissue, comprising:

3           (a) a handpiece having a tool supporting end, and a battery  
4 receiving end with an alignment post extending therefrom, the  
5 battery receiving end of the handpiece also having a set of  
6 electrical contact elements arranged in generally concentric  
7 relation to the alignment post;

8           (b) a disposable battery having an attachment end with a  
9 central opening therein, and a set of mating electrical contact  
10 elements arranged in a generally circular configuration  
11 concentric to the central opening therein;

12           (c) the central opening in the disposable battery being  
13 adapted to receive the alignment post of the handpiece in a  
14 partially inserted position so as to establish a pre-attachment  
15 alignment thereof;

16           (d) the sets of mating contacts being adapted to come into a  
17 mutually concentric relation in response to a further insertion  
18 of the alignment post into the central opening; and

19           (e) the sets of contacts being adapted to then become  
20 lockingly and conductively interengaged upon rotation of the  
21 battery pack relative to the handpiece.

12. The apparatus of Claim 11 wherein the chemistry of the  
disposable battery is based upon lithium/manganese dioxide, and  
which further includes a sterile package containing the  
disposable battery.

1           13. A surgical handpiece for performing a cutting, shaping,  
2 or drilling operation on bone or hard tissue, comprising:

3           (a) a tool supporting end;

4           (b) an electric motor for driving a tool;

5           (c) a battery receiving end having a flat end surface with a  
6 defined central axis, and a circumferentially arranged set of  
7 electrical contact elements concentric to the axis;

8           (d) the flat end surface being adapted for abutting  
9 rotatable engagement by the electrical contact elements of a  
10 battery pack and to then provide a spring-supported snap action  
11 in response to rotation of the contact elements of the battery  
12 pack into a predetermined locking position; and

13           (e) the battery receiving end of the handpiece also having  
14 a non-circular external cross-sectional configuration to indicate  
15 to the hand of an operator that correct rotational alignment of  
16 the battery pack has been achieved.

14. A surgical handpiece as in Claim 13 wherein the central  
axis of the flat end surface is defined by an alignment post  
extending therefrom.

15. A surgical handpiece as in Claim 13 wherein the flat  
end surface at one point on its circumference has a stop pin for  
limiting the rotation of the battery pack relative to the  
handpiece.

16. A surgical handpiece as in Claim 13 wherein the electrical contact elements are made as spring members.

1 17. A disposable battery pack for attachment to the  
2 handpiece of a surgical tool, comprising:  
3 a housing;  
4 internal primary batteries within the housing;  
5 the housing having a forward attachment end with a flat end  
6 surface adapted for abutting engagement with the end surface of  
7 the handpiece while yet allowing rotation of the battery pack  
8 relative to the handpiece;  
9 the flat end surface having means defining a longitudinal  
10 axis; and  
11 a set of electrical contact elements on the flat end surface  
12 concentric to the axis.

18. A disposable battery pack as in Claim 17 whose forward end has projecting flanges that are circumferentially unsymmetrical, so as to restrict its rotational position relative to the handpiece upon engagement therewith.

19. A disposable battery pack as in Claim 17 wherein the flat end surface has a central opening therein which defines the longitudinal axis, and which is adapted to insertably receive an alignment post of the handpiece.

20. A disposable battery pack as in Claim 17 whose chemistry is based upon lithium/manganese dioxide.

21. A surgical tool with detachable battery wherein the battery contains lithium/managanese dioxide, and is disposable after use into non-hazardous waste.

1        22. A surgical method comprising:  
2        selecting a handpiece adapted for removable attachment of a  
3 battery pack thereto;  
4        selecting a packaged and pre-sterilized battery pack  
5 containing primary batteries whose chemistry is based upon  
6 lithium/manganese dioxide;  
7        attaching the battery pack to the handpiece to provide  
8 electrical energy for its operation;  
9        conducting a surgical procedure utilizing the handpiece; and  
10       then disposing of the battery pack into non-hazardous waste.

23. The surgical method of Claim 22 wherein the selected surgical handpiece is a compact device containing a tool member, a brushless DC motor for moving the tool member, and a manually operated trigger for activating motor control operations.



24. The surgical method of Claim 22 wherein the selected surgical handpiece and the selected disposable battery pack have cooperating means for limiting the rotational position of the battery pack relative to the handpiece prior to their mutual engagement.

1           25. A surgical method of performing a cutting, shaping, or  
2 drilling operation on bone or hard tissue, comprising steps of:

3           (a) selecting a handpiece having a tool supporting end, and  
4 also having a battery receiving end with a set of electrical  
5 contact elements thereon;

6           (b) selecting a sterile package containing a disposable  
7 battery whose chemistry is based upon lithium/manganese dioxide,  
8 the battery having an attachment end with a set of mating  
9 electrical contact elements thereon;

10          (c) removing the disposable battery from the sterile  
11 package;

12          (d) conductively interengaging the sets of contact elements  
13 so as to provide energy for the handpiece;

14          (e) utilizing the tool to perform a surgical procedure; and

15          (f) thereafter detaching and disposing of the disposable  
16 battery into non-hazardous waste.

1           26. The method of Claim 24 wherein the handpiece and the  
2 battery are selected to have sets of contacts which are adapted  
3 to become lockingly and conductively interengaged upon rotation  
4 of the battery pack relative to the handpiece, in a manner that  
5 achieves correct alignment of the parts and also ensures stable  
6 mechanical attachment and support during the surgical procedure.

1           27. A surgical method comprising the steps of:

2           (a) selecting a handpiece having a tool supporting end, and  
3 a battery receiving end with electrical contact elements thereon;

4           (b) selecting a disposable battery having an attachment end  
5 with electrical contacts adapted to engage the tool contacts;

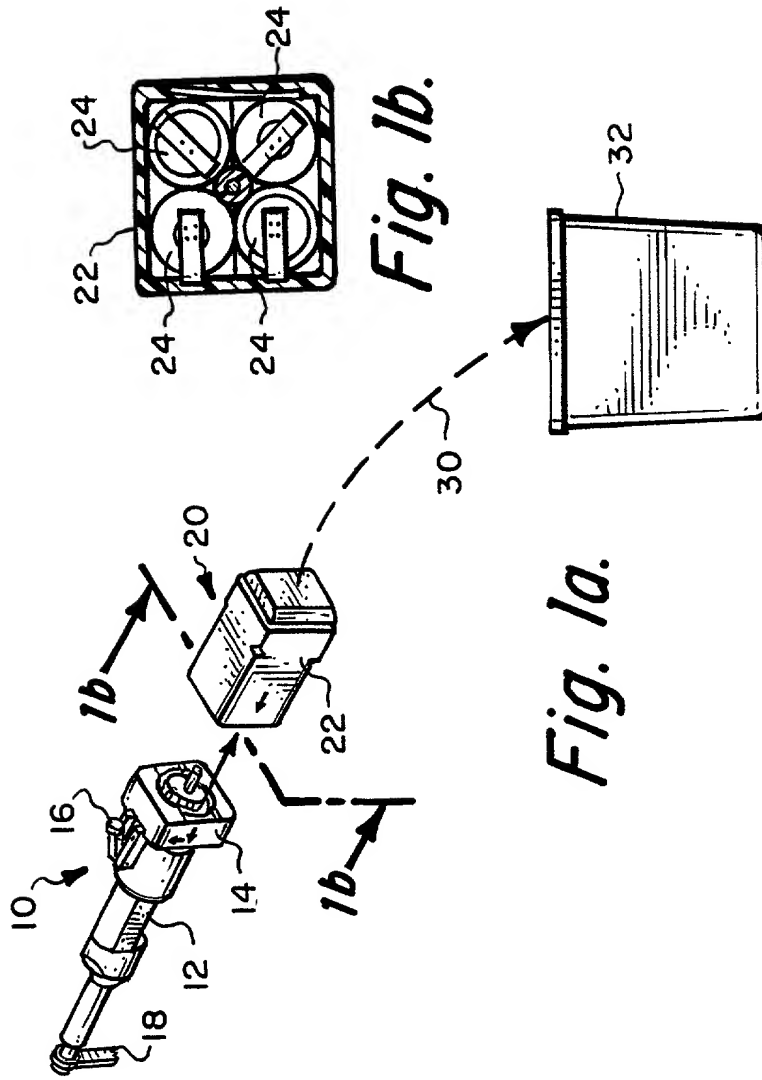
6           (c) rotatably locking the battery to the handpiece to  
7 conductively interengage the sets of contact elements, while  
8 concurrently creating a positive indication by at least one of  
9 sight, sound, and touch that a predetermined locked position has  
10 been achieved; and

11           (d) thereafter utilizing the tool to perform a cutting,  
12 shaping, or drilling surgical procedure on bone or hard tissue.

28. The method of Claim 27 wherein the battery is selected  
to have its chemistry based upon lithium/manganese dioxide.

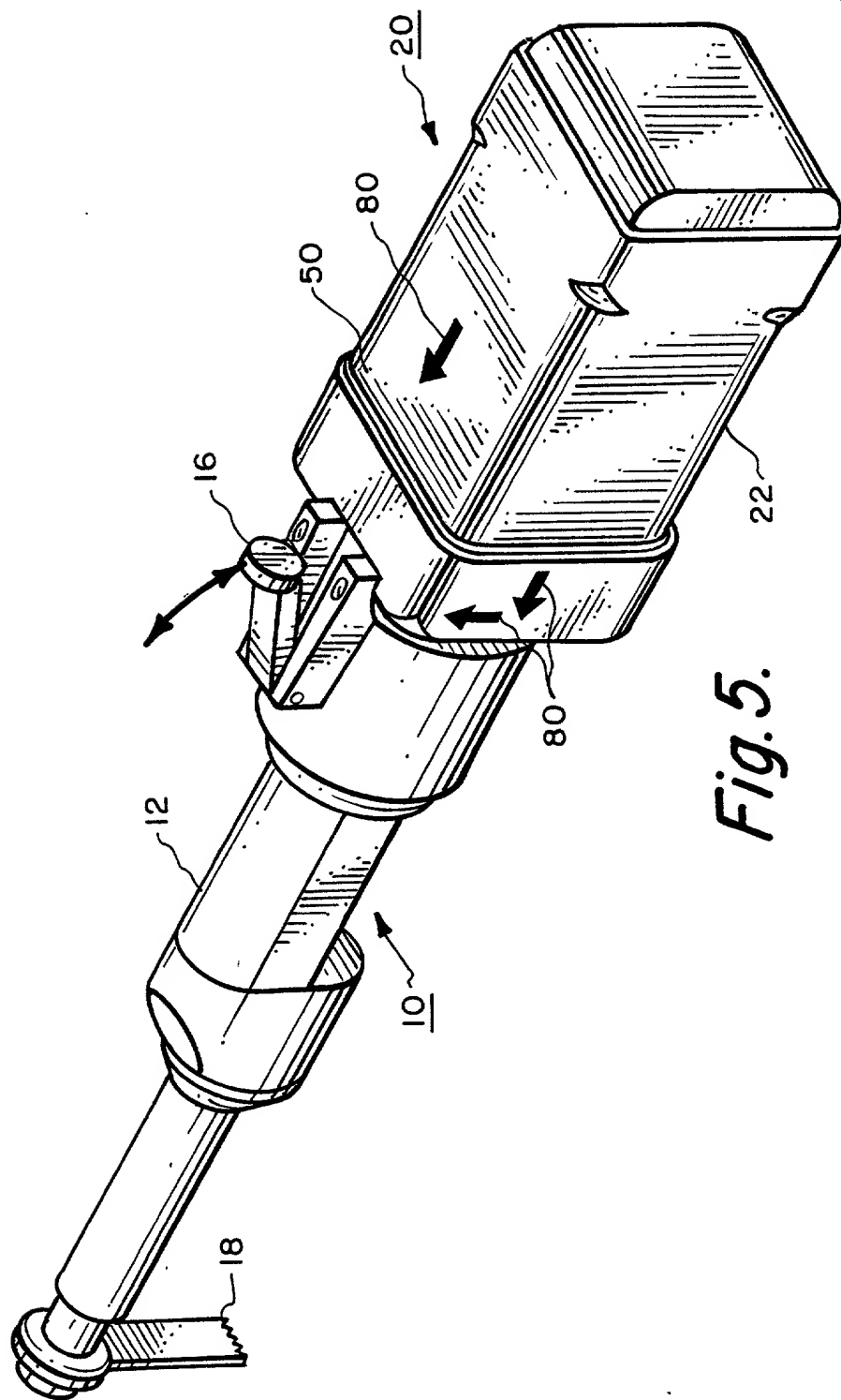
1 ABSTRACT

2 A surgical procedure is disclosed utilizing a cordless  
3 surgical handpiece powered from a sterile battery pack that  
4 contains a battery in condition for immediate use without further  
5 charging or sterilization. The battery chemistry is based upon  
6 lithium/manganese dioxide, and the battery after a single use may  
7 be disposed of into non-hazardous waste. The compact surgical  
8 handpiece has a brushless DC motor and a manually operated  
9 external trigger for activating and controlling the motor  
10 operations. Interengaging sets of contacts on the handpiece and  
11 battery are adapted to become lockingly and conductively  
12 interengaged upon rotation of the battery pack relative to the  
13 handpiece, in a manner that rapidly achieves correct alignment of  
14 the parts and also ensures stable mechanical attachment and  
15 support during the surgical procedure.

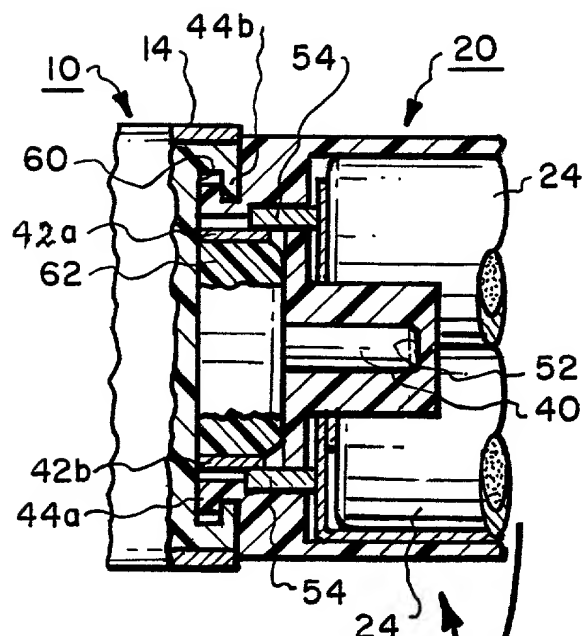
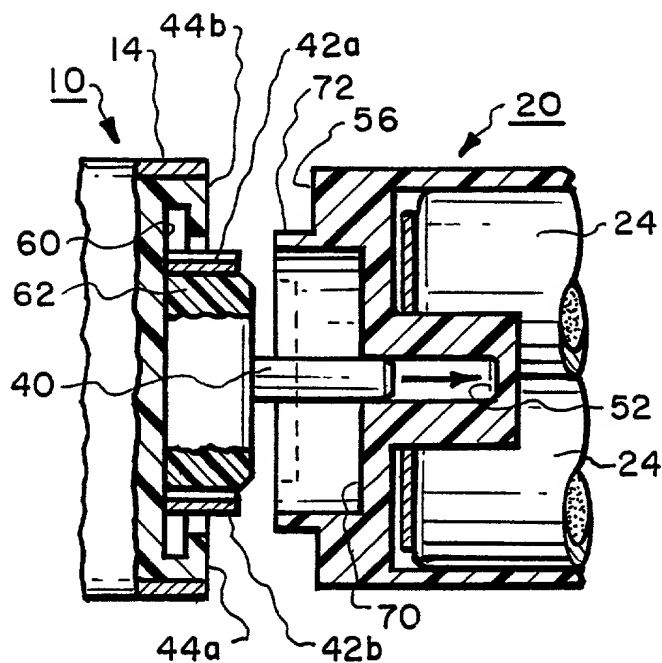
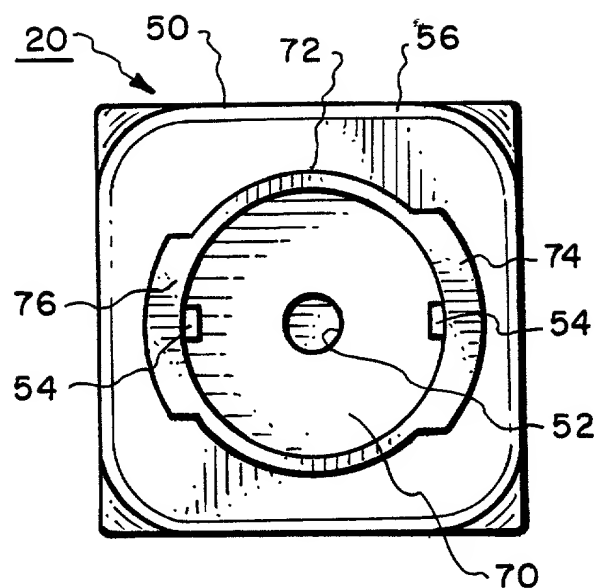
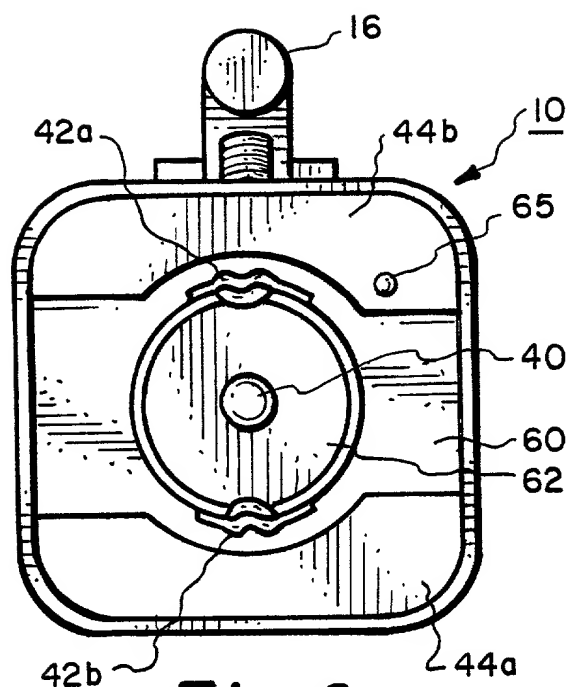




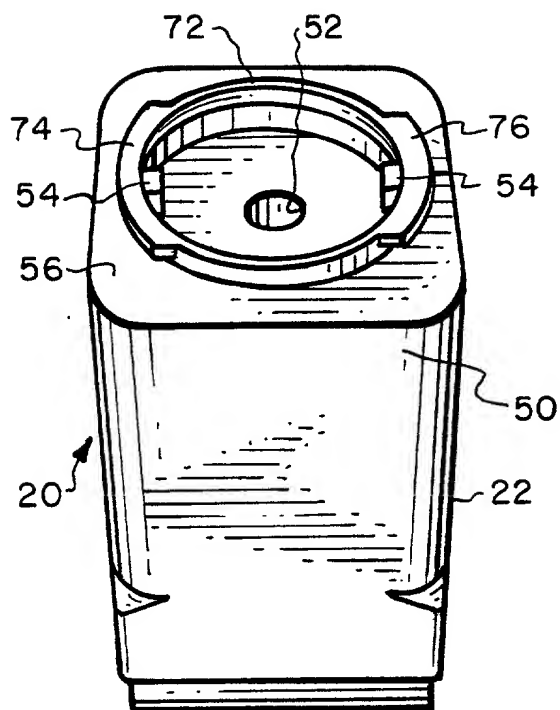




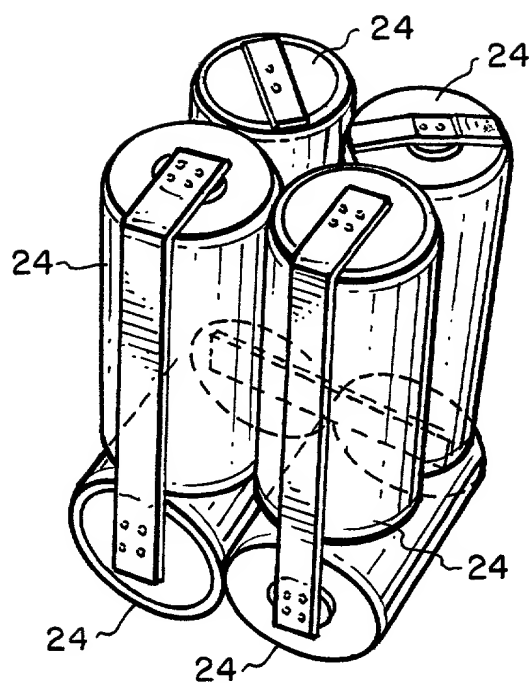
**Fig. 5.**



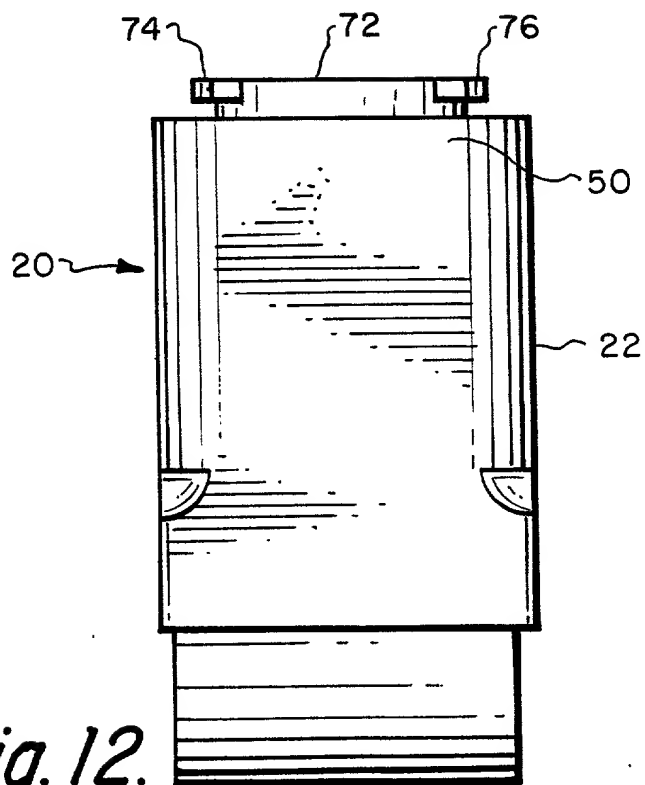




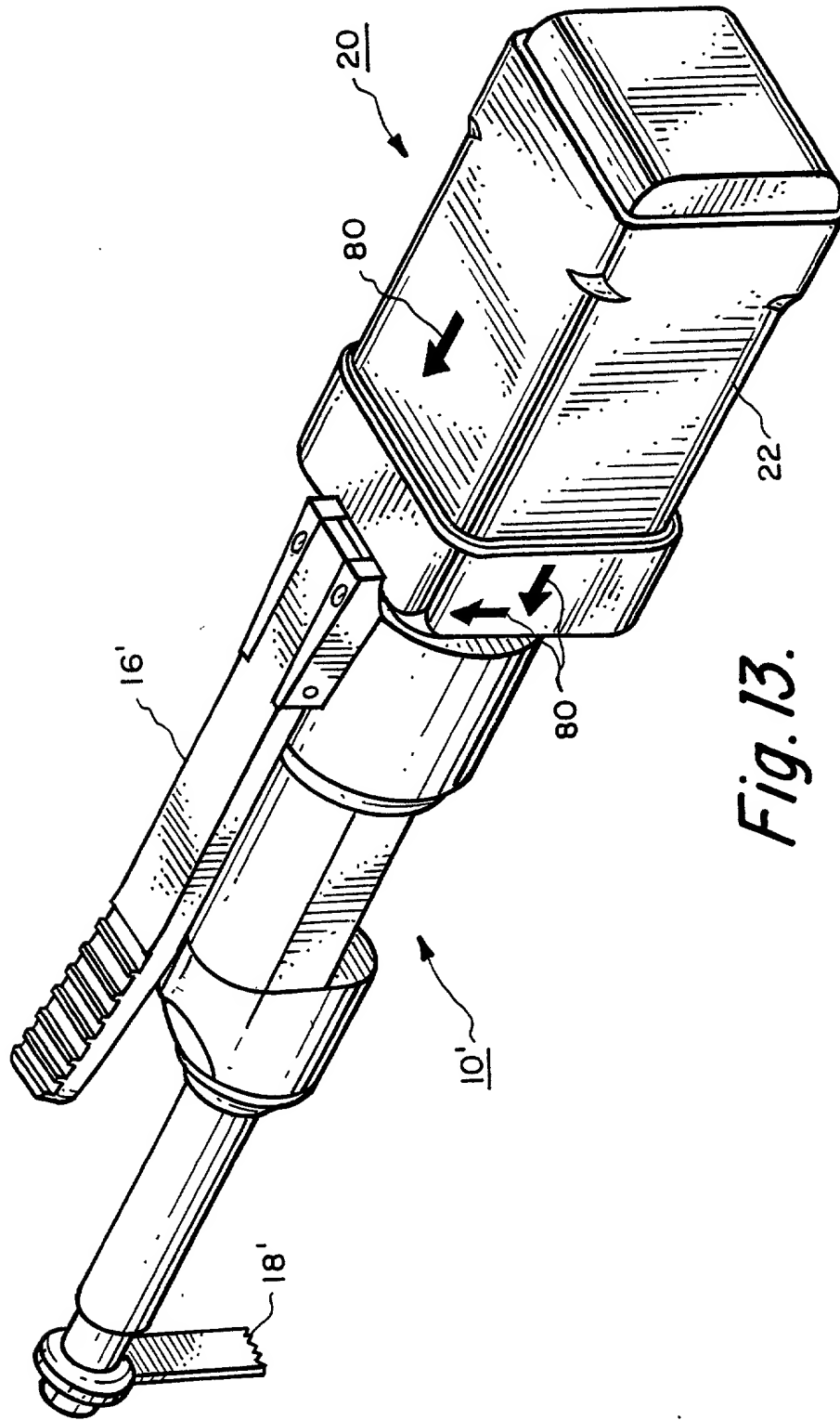
*Fig. 10.*



*Fig. 11.*



*Fig. 12.*



**Fig. 13.**

EXPRESS MAIL CERTIFICATE

Date 9/23/00 Label No EL003316715US

I hereby certify that, on the date indicated above I deposited this paper or fee with the U.S. Postal Service and that it was addressed for delivery to the Commissioner of Patents & Trademarks, Washington, DC 20231 by "Express Mail Post Office to Addressee" service.

Name (Print) Mei Kyle Signature Mei Kyle

FILE NO.: 135555-0262

**DECLARATION  
AND POWER OF ATTORNEY  
Continuing Application**

As a below named inventor, I declare that the information given herein is true, that I believe that I am the original, first and sole inventor if only one name is listed at 1 below, or a joint inventor if plural inventors are named below, of the invention entitled:

**CORDLESS SURGICAL HANDPIECE WITH DISPOSABLE BATTERY; AND METHOD**

which is described and claimed in:

☒ the attached specification

that I do not know and do not believe that the same was ever known or used in the United States of America before my or our invention thereof or patented or described in any printed publication in any country before my or our invention thereof, or more than one year prior to this application, or in public use or on sale in the United States of America more than one year prior to this application, that the invention has not been patented or made the subject of an inventor's certificate issued before the date of this application in any country foreign to the United States of America on an application filed by me or my legal representatives or assigns more than twelve months prior to this application, that I acknowledge my duty to disclose information of which I am aware which is material to the examination of this application in accordance with 37 CFR §1.56(a), and that no application for patent or inventor's certificate on this invention has been filed by me or my legal representatives or assigns in any country foreign to the United States of America except as identified below. I hereby state that I have reviewed and understand the contents of the above-identified specification, including the claims, as amended by any amendment referred to above.

**FOREIGN APPLICATION(S), IF ANY, FILED WITHIN 12 MONTHS  
PRIOR TO THE FILING DATE OF THIS APPLICATION**

None.

**ALL FOREIGN APPLICATIONS, IF ANY, FILED MORE THAN 12 MONTHS  
PRIOR TO THE FILING DATE OF THIS APPLICATION**

None.

**CLAIM FOR BENEFIT OF PRIOR U.S. APPLICATION  
UNDER 35 U.S.C. 120:**

I hereby claim the benefit under 35 U.S.C. 120 of any United States application(s), or 365(c) of any PCT international application designating the United States of America, listed below and, insofar as the subject matter of each of the claims of this application is not disclosed in the prior United States or PCT International application in the manner provided by the first paragraph of 35 U.S.C. 112, I acknowledge the duty to disclose information which is material to patentability as defined in 37 CFR 1.56 which became available between the filing date of the prior application and the national or PCT international filing date of this application.

Co-pending application Serial No. 09/343,643 filed July 8, 1999, which in turn claimed the benefit of Provisional Application Serial No. 60/112,678 filed December 12, 1998.

**POWER OF ATTORNEY:**

As a named inventor, I hereby appoint the following attorney(s) and/or agents(s) to prosecute this application *and all corresponding foreign applications* and transact all business in the Patent and Trademark office connected therewith: Gene W. Arant (Reg. No. 17,936) and Larry D. Baker (Reg. No. 39,593), of counsel, of the Law Offices of Gene W. Arant, located at 674 County Square Drive, Suite 205, Ventura, California, 93003-5452.

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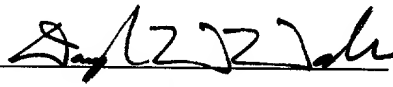
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I further declare that all statements made herein of my own knowledge are true and that all statements made on information and belief are believed to be true; and further that these statements were made with the knowledge that willful false statements and the like so made are punishable by fine or imprisonment, or both, under Section 1001 of Title 18 of the United States Code, and that such willful false statements may jeopardize the validity of the application or any patent issuing thereon.

SIGNATURE OF INVENTOR 1:  DATED: 9/20/00

SIGNATURE OF INVENTOR 2: Frank M. Ordaz DATED: 9/19/00

SIGNATURE OF INVENTOR 3: M. Terry Olson DATED: 9-19-00